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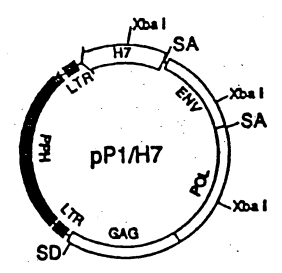
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(57) Abstract

This invention relates to a method of immunizing a vertebrate, comprising introducing into the vertebrate a DNA transcription unit which comprises DNA encoding a desired antigen or antigens. The uptake of the DNA transcription unit by a host vertebrate results in the expression of the desired antigen or antigens, thereby eliciting humoral or cell-mediated immune responses or both humoral and cell-mediated responses. The elicited humoral and cell-mediated immune response can provide protection against infection by pathogenic agents, provide an anti-tumor response, or provide contraception. The host can be any vertebrate, avian or mammal, including humans.

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IMMUNIZATION BY INOCULATION OF DNA TRANSCRIPTION UNIT

Background of the Invention

Vaccination with inactivated or attenuated organisms

or their products has been shown to be an effective method for increasing host resistance and ultimately has led to the eradication of certain common and serious infectious diseases. The use of vaccines is based on the stimulation of specific immune response within a host or the transfer of preformed antibodies. The prevention of certain diseases, such as poliomyelitis, by vaccines represents one of immunologies greatest triumphs.

Effective vaccines have been developed for relatively few of the infectious agents that cause disease in domestic animals and man. This reflects technical problems associated with the growth and attenuation of virulent strains of pathogens. Recently effort has been placed on the development of subunit vaccines (vaccines that present only selected antigens from a pathogen to the host). Subunit vaccines have the potential for achieving high levels of protection in the virtual absence of side effects. Subunit vaccines also offer the opportunity for the development of vaccines that are stable, easy to administer, and sufficiently cost-effective for widespread distribution.

Summary of the Invention

This invention relates to a method of immunizing an individual, comprising introducing into the individual a DNA transcription unit which comprises DNA encoding a desired antigen or antigens. The uptake of the DNA transcription unit by host cells results in the expression of the desired antigen or antigens, thereby eliciting

humoral or cell-mediated immune responses or both humoral and cell-mediated responses. The elicited humoral and cell-mediated immune response can provide protection against infection by pathogenic agents, provide an antitumor response, or provide contraception. The host can be any vertebrate, avian or mammal, including humans.

The present invention relates in a particular embodiment to a method of immunizing an individual by contacting a mucosal surface in the individual with a DNA transcription unit capable of expressing a desired antigen or antigen.

The DNA transcription unit introduced by the present method can be used to express any antigen encoded by an infectious agent, such as a virus, a bacterium, a fungus, or a parasite, as well as antigenic fragments and peptides that have been experimentally determined to be effective in immunizing an individual against infection by a pathogenic agent. As stated above, DNA transcription units can also be used for contraceptive purposes or for anti-cancer therapy.

The desired antigen to be expressed can be designed so as to give internal, surface, secreted, or budding and assembled forms of the antigens being used as immunogens.

There are numerous advantages for the use of DNA for immunizations. For example, immunization can be accomplished for any antigen encoded by DNA. Furthermore, the DNA encoded antigens are expressed as "pure" antigens in their native states and have undergone normal host cell modifications. Also, DNA is easily and inexpensively manipulated and is stable as a dry product or in solution over a wide range of temperatures. Thus, this technology is valuable for the development of highly effective subunit vaccines.

Brief Description of the Drawings

Figure 1 is an illustration of a bacterial plasmid containing a DNA transcription unit (referred to as pP1/H7) comprising an influenza virus hemagglutinin type 7 (H7) gene expressed by a replication competent retroviral vector.

Figure 2 is an illustration of a bacterial plasmid containing a DNA transcription unit (p188) comprising an influenza virus hemagglutinin type 7 (H7) gene expressed by a replication defective retroviral vector.

Figure 3 is an illustration of a bacterial plasmid comprising a retroviral vector (pRCAS) with no H7 insert, used as a control.

Figure 4A is a schematic representation of the nonretroviral vector comprising the influenza virus antigen DNA transcription unit encoding subtype H7 hemagglutinin.

Figure 4B is a schematic representation of the nonretroviral vector comprising the influenza virus antigen DNA transcription unit encoding subtype H1 hemagglutinin.

Figure 4C is a schematic representation of the nonretroviral vector comprising a control DNA transcription unit, encoding no influenza virus antigens.

Figure 5 is a bar graph depicting the maximum median weight loss for DNA-vaccinated mice in experiment 4, Table 7.

Detailed Description of the Invention

This invention relates to a method of immunizing
vertebrates, particularly mammals, including humans,
against a pathogen, or infectious agent, thereby eliciting
humoral and/or cell-mediated immune responses which limit
the spread or growth of the infectious agent and result in

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protection against subsequent challenge by the pathogen or infectious agent.

The term "immunizing" refers herein to the production of an immune response in a vertebrate which protects

(partially or totally) from the manifestations of infection (i.e., disease) caused by an infectious agent. That is, a vertebrate immunized by the present invention will not be infected or will be infected to a lesser extent than would occur without immunization.

A DNA transcription unit is a polynucleotide sequence which includes at least two components: antigen-encoding DNA and transcriptional promoter elements. A DNA transcription unit may optionally include additional sequences, such as: enhancer elements, splicing signals, termination and polyadenylation signals, viral replicons and bacterial plasmid sequences.

The DNA transcription unit can be produced by a number of known methods. For example, using known methods, DNA encoding the desired antigen can be inserted into an expression vector to construct the DNA transcription unit. See Maniatis et al., Molecular Cloning. A Laboratory Manual, 2d, Cold Spring Harbor Laboratory Press (1989).

The DNA transcription unit can be administered to an

25 individual, or inoculated, in the presence of adjuvants or
other substances that have the capability of promoting DNA
uptake or recruiting immune system cells to the site of
the inoculation. It should be understood that the DNA
transcription unit itself will be expressed by host cell

30 factors.

The "desired antigen" can be any antigen expressed by an infectious agent or any antigen that has been determined to be capable of eliciting a protective response against an infectious agent. These antigens may or may not be structural components of the infectious

agent. The encoded antigens can be translation products or polypeptides. The polypeptides can be of various lengths. They can undergo normal host cell modifications such as glycosylation, myristoylation or phosphorylation.

In addition, they can be designed to undergo intracellular, extracellular or cell-surface expression.

Furthermore, they can be designed to undergo assembly and release from cells.

Potential pathogens for which the DNA transcription
10 unit can be used include DNA encoding antigens derived
from any virus, chlamydia, mycoplasma, bacteria, parasite
or fungi. Viruses include the herpesviruses
orthomyxoviruses, rhinoviruses, picornaviruses,
adenoviruses, paramyxoviruses, coronaviruses,
rhabdoviruses, togaviruses, flaviviruses, bunyaviruses,
rubella virus, reovirus, hepadna viruses and retroviruses
including human immunodeficiency virus.

rubella virus, reovirus, hepadna viruses and retroviruses including human immunodeficiency virus. Bacteria include mycobacteria, spirochetes, rickettsias, chlamydia, and mycoplasma. Fungi include yeasts and molds. Parasites include malaria. It is to be understood that this list

does not include all potential pathogens against which a protective immune response can be generated according to the methods became described.

the methods herein described.

An individual can be inoculated through any

parenteral route. For example, an individual can be
inoculated by intranasal, intravenous, intraperitoneal,
intradermal, subcutaneous or intramuscular methods. In a
particular embodiment of the present invention, an
individual is vaccinated by contacting a mucosal surface

on the individual with the desired DNA transcription unit
in a physiologically compatible medium. The DNA
transcription unit can be administered to a mucosal
surface by a variety of methods, including DNA-containing
nose-drops, inhalants and suppositories.

Any appropriate physiologically compatible medium, such as saline, is suitable for introducing the DNA transcription unit into an individual.

The following Examples describe vaccination trials using direct DNA inoculations designed for use in both avian and murine influenza virus models. Both of these models afford rapid assays for protective immunizations against lethal challenges, wherein challenge of an unimmunized animal causes death within 1-2 weeks.

Immunization as described herein has been
accomplished with DNA transcription units (i.e., vectors)
that express an influenza virus hemagglutinin
glycoprotein. This protein mediates adsorption and
penetration of virus and is a major target for
neutralizing antibodies. Influenza virus hemagglutinin
proteins have 14 different serological subtypes. In the
avian model, DNA expression vectors for the H7 subtype
(comprising a DNA transcription unit encoding the H7
subtype hemagglutinin) have been used to provide
protection against challenge with an H7N7 virus. In the
murine model, a DNA transcription unit expressing the H1
hemagglutinin was used to immunize against an H1N1 virus.

<u>Example 1 - Immunization of Chickens Against Influenza</u> Virus

25 Procedure:

A DNA transcription unit referred to as pP1/H7 (Fig. 1), encoding a replication competent avian leukosis virus expressing the influenza virus hemagglutinin type 7 (H7) gene was constructed as described in Hunt et al., J. of

30 Virology, 62(8):3014-3019 (1988). DNA unit p188 (Fig. 2) encoding a replication defective derivative of pP1/H7 that expresses H7 but is defective for the avian virus vector polymerase and envelope proteins was constructed by deleting an XbaI fragment from pP1/H7. DNA unit pRCAS

(Fig. 3), encoding the avian leukosis virus vector, with no influenza virus insert, was constructed as described in Hughes et al., J. of Virology, 61:3004 (1987). DNA units were diluted in saline at a concentration of 100 μ g per 0.2 ml for inoculation.

To test the ability of the inoculated DNA to protect against a lethal influenza virus challenge, groups of three-week old chicks were inoculated with pP1/H7, p188, or pRCAS DNA. Specific pathogen free chicks that are maintained as an avian-leukosis virus-free flock (SPAFAS, 10 Norwich, CT) were used for inoculations. Each chick received 100 μ g of DNA (~1x10¹³ molecules) intravenously (iv), 100 μ g intraperitoneally (ip), and 100 μ g subcutaneously (sc). Four weeks later chicks were bled and boosted with 300 μg of DNA (100 μg iv, 100 μg ip, and 100 μ g sc). At one week post-boost, chicks were bled and challenged by the nares with 100 lethal doses (1x104 egg infectious doses) of a highly pathogenic type H7 avian influenza virus, A/Chicken/Victoria/1/85 (H7N7) 20 (Ck/Vic/85). The chickens were observed daily for ten days for signs of disease. One and one half weeks after challenge, sera were obtained from surviving birds. as well as the pre- and post-boost sera were used for analyses for hemagglutination inhibiting antibodies (HI).

Sera were analyzed in microtiter plates with receptor-destroying enzyme-treated sera as described by Palmer et al., Advanced Laboratory Techniques for Influenza Diagnosis, p. 51-52, Immunology series no. 6, U.S. Department of Health, Education, and Welfare, 30 Washington, D.C. (1975).

Results:

The H7-expressing DNA transcription units protected each of the chickens inoculated with pP1/H7 or p188 (Table 1). In contrast, inoculation with the control DNA, pRCAS,

failed to protect the chickens against lethal virus challenge. The birds in the control group started to show signs of disease on the second day post-challenge. By the third day, three of the six control birds had died and all control birds were dead by the fifth day. The birds inoculated with hemagglutinin-expressing DNAs showed no signs of disease. By one and one half weeks post challenge both of these groups had developed high levels of HI antibody.

10 Example 2 - Immunization Against Influenza Virus is Reproducible

To assess the reproducibility of the protection elicited by immunization with the replication-defective H7-expressing DNA, the experiment described in Example 1 15 was repeated three times using only p188 and pRCAS DNAs for inoculations. The results of the repeat experiments confirmed that the H7-expressing pl88 DNA could afford protection against a lethal challenge (Table 2). contrast to the first experiment, in which all of the p188-inoculated chickens survived the lethal challenge, immunizations in the 2nd, 3rd, and 4th experiments achieved only partial protection with from 28% to 83% of the vaccinated birds surviving. Further, in contrast to the first experiment in which vaccinated birds showed no signs of disease, most of the survivors of the repeat experiments showed transient signs of post-challenge sickness. As in the first experiment, the control DNA did not provide protection. Summing the results of the 4 experiments, 28 out of 56 pl88-vaccinated birds survived 30 whereas only 1 of 55 control DNA-inoculated birds survived. Thus, despite the variability, significant immunization was achieved.

Example 3 - Immunization can be Accomplished by Several Different Routes of Inoculation

Procedure:

The DNA encoding p188-H7 and control DNA were tested 5 again for the ability to protect against a lethal influenza virus challenge. This experiment included a group that was vaccinated and boosted by three routes of inoculation (i.e., ip, iv and sc), a group that was vaccinated by the same three routes but did not receive a 10 boost, small groups that were vaccinated and boosted by only one route of inoculation and a control group treated with the anti-influenza virus drug, amantadine-HCL. last group was included to allow the comparison of antibody responses to the challenge virus in vaccinated and unvaccinated chickens. 15 The amantadine-treated birds were given 0.01% amantadine in their drinking water beginning 8 hours after challenge and were also injected ip with 1.0 ml of 0.1% amantadine 24 and 48 hours after challenge.

20 Results:

The results of this experiment confirmed that the replication defective H7-expressing DNA (p188) could afford protection against a lethal virus challenge (Table 3). The p188 immunized birds showed transient signs of sickness following the challenge. As in the previous experiments, the control DNA did not provide protection. All of the 5 amantadine-treated control birds developed disease. Four of these survived the challenge, providing sera that could be used to compare the time course and specificity of anti-influenza virus responses in immunized and non-immunized chickens (see Example 5 below).

Example 4 - Immunization can be Accomplished by Several Different Routes of Inoculation

Procedure:

A third experiment was initiated to increase the

numbers of birds in the test groups and to further
evaluate the efficacy of different routes of immunization.
In this experiment 12 chicks were inoculated with 100 μg
p188 by the iv, ip, and sc routes, 8 chicks were
inoculated iv-only and 8 ip-only. For controls, 12 chicks
were inoculated with pRCAS and 12 chicks were not
inoculated. All immunizations were followed by a boost
four weeks after the initial inoculation. The boosts used
the same DNA dose and sites of inoculation as the
vaccinations. The control and immunized animals were

challenged with ck/vic/85 1-2 weeks after the boost, with
high challenge doses used in order to achieve essentially
100% killing within 1-2 weeks.

Results:

The results again demonstrated protection by p188

20 (Table 4). Eight of the 12 p188 immunized birds survived, whereas all 12 of the control pRCAS chickens died. The twelve birds in the untreated control group also had no survivors. Six out of the 8 chickens inoculated iv-only with p188 survived whereas none of the 8 chickens

25 inoculated ip-only survived.

Example 5 - Analysis of Antibody Response to Challenge Virus in Vaccinated and Unvaccinated Animals

Procedure:

To allow the comparison of antibody responses to the 30 chall nge virus in vaccinated and unvaccinated chickens, experiment 2 from Example 2 (Table 2) included a non-vaccinated group rescued with the anti-influenza A virus drug, amantadine-HCL (Table 2) (Webster, R.G., et al., J.

<u>Virol.</u> 55:173-176 (1985)). All of the 5 amantadinetreated birds developed disease. Four of these survived, providing sera that could be used to compare antibody responses in immunized and non-immunized chickens (Table 6).

Sera from p188 inoculated and amantadine treated birds in the second experiment were analyzed for the time course of antibody responses to H7 and to other influenza virus proteins (Table 6). Antibody responses to H7 were 10 quantitated using hemagglutination inhibition as well as virus neutralization and enzyme-linked immunosorbent assays (ELISA) for antibody. Neutralizing antibody was determined in chick embryo fibroblast cultures with 200 ${\tt TCID}_{\tt 50}$ of virus using cytopathology and hemagglutinin for detection of virus replication.

Results:

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Analysis of the antibody responses in vaccinated and amantadine-rescued birds revealed that the p188inoculations had primed an antibody response to H7 (Table 6). As in experiment 1 (Table 1), DNA vaccination - 20 and boost induced only low titers of antibody to H7. However, within one week of challenge, the DNA-immunized group had high titers of HI and neutralizing activity for These titers underwent little (if any) increase over the next week. Furthermore, most of the post-challenge antibody in the vaccinated birds was directed against H7. This specificity was shown by comparing ELISA antibody titers to H7 virus (the immunizing hemagglutinin type) and H5 virus (a hemagglutinin type to which the birds had not been exposed). The post-challenge sera contained 20-times higher titers of ELISA antibody for the H7 than the H5 virus (Table 6). By contrast, in the amantadine-rescued group, antibodies did not appear until two weeks postchallenge. Most of this response was not H7-specific.

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This was demonstrated by the post-challenge sera from the amantadine-rescued birds which had comparable titers of ELISA antibody for the H5 and the H7 influenza viruses (Table 6).

5 Example 6 - Immunization of Chickens and Mice Using a Nonretroviral Transcription Unit

Procedure

This experiment was performed in order to demonstrate that DNA transcription units devoid of retroviral DNA could be successfully employed to generate a protective immune response in both chickens and mice according to the methods herein described. The vectors used in this experiment to vaccinate chicken and mice are shown in Figure 4A-4C. Figure 4A is a schematic representation of pCMV-H7, a plasmid capable of expressing the influenza virus H7 subtype hemagglutinin under the transcription control of a cytomegalovirus (CMV) immediate early promoter. 4B is a schematic showing pCMV-H1, a plasmid capable of expressing the influenza virus H1 subtype hemagglutinin under the control of a CMV immediate early promoter. This is the DNA transcription unit used in the mouse experiments. Figure 4C shows pCMV, a control plasmid which is not capable of expressing influenza antigens. These plasmids are derivatives of the pBC12/CMV vector of Dr. Brian Cullen, Duke University, Durham, North Carolina.

In the chicken and mouse experiments using pCMV-H7 and pCMV-H1 DNAs (the nonretroviral-based DNA transcription units) to generate immune responses, 100 μ g of DNA was inoculated intravenously, intraperitoneally, and intramuscularly. All vaccinations were followed by a boost 4 weeks later. The boosts used the same DNA dose and sites of

inoculation as the vaccinations. Challenge was 1-2 weeks after the boost, with high challenge doses being used so as to achieve essentially 100% killing within 1-2 weeks.

5 Results:

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In five chicken trials using a nonretrovirus-based vector for vaccination (pCMV-H7) (Figure 4A), approximately 60% of the chickens were protected. In one mouse trial, six out of six vaccinated mice and only one out of six control mice survived. Thus, considerable protection has been achieved using nonretroviral DNA expression vectors (containing DNA transcription units encoding viral antigens) to vaccinate animals. See, for example, Table 5.

In the chicken experiments, protective responses were associated with the rapid appearance of H7-specific antibodies after challenge (Robinson et al., 1993). Sera contained low to undetectable levels of anti-H7 antibodies after vaccination and boost. The first mouse experiment was similar to the chicken experiments in that inoculated mice also had low titers of anti-hemagglutinin activity before challenge. However, as in the chicken experiments, high titers of antibody appeared after challenge. The vast majority of this antibody was IgG.

Example 7 - Immunization of Mice by Vaccination with a Nonretroviral Transcription Unit: Analysis of Various Routes of Inoculation

Procedure:

A DNA transcription unit referred to as pCMV-H1 (described in Figur 4B) was successfully used to immunize mice against a lethal challenge with mouse adapted A/PR/8/34 H1N1 influenza virus. This

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transcription unit encodes an influenza type H1 hemagglutinin under the transcription regulation of a CMV immediate early promoter. The H1 influenza virus hemagglutinin gene used in this construct is described in more detail in Winters et al., Nature 292:72 (1981).

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The first experiment was conducted by inoculation of 6-8 week old Balb/C mice with 100 μ g of pCMV-H1 DNA by each of three routes; iv ip and im. The second, third and fourth experiments each included one group of mice inoculated iv, ip and im, as well as additional groups representing different routes of inoculation (data summarized in Table 7 and Figure 5.

The numbers in Table 7 represent the number of surviving mice/number of inoculated mice. The routes 15 of inoculation (iv, intravenous; ip, intraperitoneal; im, intramuscular; sc., subcutaneous; in, intranasal; id, intradermal) for each trial are indicated. In most instances, 100 µg of DNA was administered per injection. Intramuscular (im) inoculations were given by injection of 100 μ g DNA in each hip muscle. 20 Intravenous (iv) inoculations were given by injection in the tail vein. Intranasal (in) administrations of DNA and challenges were done on Metofane-anesthetized animals (Pitman-Moore) (these animals inhale deeply).

Intradermal (id) inoculations were done in the foot pad using only 50 μg of DNA. The control groups in experiments 2 and 3 received saline. The controls for experiment 1 received control DNA (vector without an insert encoding the antigen) administered iv, ip and im. The control group in experiment 4 received control DNA im, in and id. Occasional mice are resistant to influenza challenge. One of the survivors in the intranasal group in experiment 2, the one survivor in the control group in experiment 1, and 1 survivor in the control group in experiment 4 were such resistant

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mice. All groups showed signs of sickness following challenge. Data on weight loss were also collected and are presented in Figure 5. The weight loss data provides a quantitative measure for the degree of sickness in the different experimental groups.

Results:

The survival data, weight loss data and initial serology data from this series of experiments indicate that many routes of inoculation can provide protective In addition, these data demonstrate that 10 immunity. intranasal inoculation (DNA nose drops administered to Metofane-anesthetized mice) can provide protective immunity to a lethal virus challenge. The method herein described may, therefore, provide means of stimulating mucosal immunity. (Table 7 and Figure 5]. 15 Finally, these data demonstrate that some routes of inoculation are more effective than others for generating a protective immune response (Table 8).

Example 8 - Antibody Responses to Challenge Virus in Animals Vaccinated with a Nonretroviral DNA Transcription Unit

Experiments analyzing the serum response in pCMV-H7-vaccinated chickens were performed as described in Example 4. pCMV-H7 immunizations primed antibody responses, with high titers of antibody to H7 appearing post-challenge (Table 9).

-16-

TABLE 1 - Protection Against Lethal H7N7 Influenza Virus with DNA Coding for H7 Hemagglutinin

			HI TITERS	3
Group	Sick/Dead/Total	Post- vaccine 4 weeks	Post- boost 1 week	Post- Challenge 1.5 weeks
pP1/H7	0/0/6	<. a	<.	864 (160-1280)
p188	0/0/6	p	<	427 (160-1280)
pRCAS	6/6/6	<	<	+

a (<.) means one of six birds had an HI titer of 10.

b (<) means that all birds had titers of less than 10.

c (+) means that all birds died.

- Reproducibility of Protection Against a Lethal H7 Virus Challenge by Immunization with an H7-expressing DNA⁸

tested)
/number
c of survivors/number tested)
of
(number
droab
challenge
of
ate

			•				
	No treatment	1	1	1	0/12	0/12	
	Amantadine	1	4/5	1	i	4/5	
	PRCAS DNA	9/0	1/5	0/32	0/12	1/55	
	p188 DNA	9/9	9/9	9/32	8/12	28/56	
Exper-	iment	r i	2	က	.4	Total	

week post boost in experiment 1 and at two weeks post boost in experiments 2, Challenge was at one Experiment 1 is the same as that presented in Table 1. 3 and 4, -, not tested.

One to two weeks Three-week-old SPAFAS chicks were inoculated with 100 μg of DNA by each of Some survivors suffered transient signs of influenza later, chickens were challenged via the nares with 100 lethal doses of Four weeks later, they were boosted by inoculation with 100 μg of DNA administered iv, ip and sc. three routes (iv, ip and sc). A/CK/V1c/85 (H7N7). virus infections.

TABLE 3 - Protection Against Lethal H7N7 Influenza Virus with DNA Coding for H7 Hemagglutinin

Group	Route of Inoculation	Boost	Sick/Dead/Total ^a
p188	ip/iv/sc	yes	6/1/6
p188	iv only	yes	1/1/2
p188	ip only	yes	0/0/2
p188	sc only	yes	2/2/2
pRCAS	ip/iv/sc	yes	5/4/5
none	иА ^Ъ	NA	
none Aman. ^c	NA	NA	5/1/5
p188	iv/ip/sc	no.	4/4/6
pRCAS	iv/ip/sc	no	6/6/6

Sick birds that survived developed only mild signs of sickness such as ruffled feathers and temporary loss of appetite.

ь

⁽NA) not applicable. (Aman.) is an abbreviation for Amantadine. C

TABLE 4 - Protection Against Lethal H7N7 Influenza Virus with DNA Coding for H7 Hemagglutinin

Group	Route of Inoculation	Boost	Sick/Dead/Total ^a
p188	iv/ip/sc	yes	6/4/12
p188	iv only	yes	2/2/8
p188	ip only	yes	8/8/8
pRCAS	iv/ip/sc	yes	12/12/12
none	NAb	. AN	12/12/12

Sick birds that survived developed only mild signs of sickness such as ruffled feathers and temporary loss of appetite.

(NA) not applicable

TABLE 5 - Protection Against a Lethal H7 Influenza Virus Challenge by Immunization with pCMV-H7 DNA.

Fate of challenge group (number of survivors/number tested)				
pcmv-H7 DNA	pcmv dna			
5/6	. 0/6			
4/6	0/6			
2/6	0/7			
4/6	1/7			
4/6	0/7			
19/30	1/33			
	of survivors/number pCMV-H7 DNA 5/6 4/6 2/6 4/6 4/6			

Immunization and boosts were the same as in Table 2. Some survivors developed transient signs of influenzarelated illness.

TABLE 6- Antibody Response in H7-Immunized and Amantadine-Treated Birds

					Antibody to CK/Vic/85 (H7N7)		Antibody to Ck/Penn/1370/83 (H5N2)	
Grp.	No.ª	Bleed	þ	HI	Neutralizing	ELISA (x10 ⁻³)	ELISA (x10 ⁻³)	
p188	. 9	1 WK	wk PBb	5 (0-10)	2 (0-10)	2 (0-10)	v	
	9	2 wk	c PB	8 (0-20	13 (0-33)	5 (0-10)	•	
	Ŋ	H XX	wk PCc	112 (80-160)	873 (33-333)	640 (100-1000)	26 (0-100)	
	ທ	2 wk	S PC	272 (80-640)	540 (33-1000)	640 (100-1000)	46	-
								-21-
None	ഗ	1 wk	r PB	٧ و		v	٧	
	Ŋ	2 vk	c PB	v				
	4	1 wk	PC 1	v	. •	v		
	4	2 WK PC	DE 1	300 (80-640)	442 (100-1000)	1000 (1000)	1000 (1000)	

Antibody titers are given as the median (range). (No.) Number of chicks in group at time of bleed.

(wk PB) means weeks post boost. o o

means all birds had titers of less than 10. means weeks post challenge. (wk PC) **⊙**

TABLE 7 - Survival Data for Four DNA Immunization Trials Using pCMV-H1 in the Murine/Influenza Virus Model

,	iv,ip,		•	-, -,	7	Ç	ָב <u>.</u>	
Control	im,	E1	u I	>	ָם י	ָ מ	<u>)</u>	
1/6	9/9							
9/0	9/9	9/6	9/9	4/6		4/6	9/0	
9/0	9/9	9/9	3/6	9/9	9/9	٠		
2/6	3/4	1/7	4/5		3/6			
3/24	21/22	18/19	13/17	10/12	9/12	4/6	9/0	

TABLE 8 - HI Antibody Titers Following Inoculation of pCMV-H1

Time of bleed	Trial	Control	iv, ip, im,	in	in	, ,	Ţ	ບ
Prebleed	4 0 E 4	v v v v	v v v v	v v v	. • • •	· v v	v v	v
4 wk PV (preboost)	4064	v v v <u>v</u>	^ ^ 4 ^	v v v	v v v	v v	v v	v
10 da PB (prechallenge)	H W M 4	v v v v	^ 4 8 ^ 0 8 ^	^ <u>^</u> 4	v v v	> 4	v v	v
4-5 da PC	H 20 E 4	v v	8 v	^ 4, 0,	v v	8	v v	v
14-19 da PC	1 2/c 4	* * * * * • • • • •	2560 640 160 640	320 320 640	320 640 640	320	640 640	640

Designations and titers are the same as those in Table 9 with the exception of: control; da, days. *One surviving mouse had a titer of 80. **Two surviving mice had titers of 320. **Two surviving mice had titers of Serology for trials reported in Table 7. Data is for pooled sera. *One surviving mouse had a titer of 80.

TABLE 9 - Antibody Responses to the H7 Challenge Virus in pCMV-H7 and pCMV-control DNA inoculated chickens

ated	ELISA (x10 ⁻³)	v v v v	, v v v v v v v v v v v v v v v v v v v	765 1000 775 1000
CMV-H7-DNA-inoculated	Neut- ralizing	~ ~ ~ ~	v v v v	33 33 108
CMV-H.	HI	N V V N	N N N N N N N N N N N N N N N N N N N	60 60 100 140
lated	ELISA (x10 ⁻³)	v v v v	v v v v	០០ភ្នំ០
Control-DNA-inoculated	Neut- ralizing	v v v v	v v v v	<u>.</u>
Cont	H	v v v v	• • • • • • • • • • • • • • • • • • •	ممِّم
	Trial	.γω4η.	0 tu 4 tu	и <u>ü</u> 4 п
	Time of bleed	4 wk PV (preboost)	1 wk PB (pre- challenge)	2 wk PC

dead. except for: PV, post vaccination and D, Its post challenge titers were HI, 80; Control birds did not receive DNA. Designations and titers are as in Table 3 '*One control bird survived in this group. Neutralizing antibody, 10; and ELISA, 100.

-25-

Equivalents

Those skilled in the art will recognize, or be able to ascertain, using no more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. These and all other such equivalents are intended to be encompassed by the following claims.

CLAIMS

- 1. A product for use in vertebrate therapy, e.g., immunization, contraception or tumor therapy, and comprising a DNA transcription unit comprising DNA encoding a desired therapeutic agent operatively linked to a promoter region.
- Use of a DNA transcription unit comprising DNA encoding a desired antigen operatively linked to a promoter region, for the manufacture of a medicament for use in vertebrate immunization by eliciting a humoral immune response, a cell-mediated immune response or both against the desired antigen.
- 3. A method of immunizing a vertebrate, said method comprising administering to a vertebrate a DNA transcription unit comprising DNA encoding a desired antigen operatively linked to a promoter region, whereby a humoral immune response, a cell-mediated immune response or both is elicited against the desired antigen.
- 20 4. Use according to Claim 2 or a method according to Claim 3 wherein the desired antigen is capable of eliciting a protective immune response against an infectious agent.
- 5. Use according to Claim 2 or Claim 4, wherein the
 medicament comprises a physiologically acceptable
 carrier and is adapted to be administered by a route
 chosen from mucosal, intranasal, intravenous,
 intramuscular, intraperitoneal, intradermal and
 subcutaneous.

- 6. The method of Claim 3 or Claim 4, wherein the DNA transcription unit, in a physiologically acceptable carrier, is administered to a vertebrate through a route of administration chosen from intranasal, intravenous, intramuscular, intraperitoneal, intradermal and subcutaneous.
- The method of Claim 3 or Claim 4, wherein the DNA transcription unit is administered to a vertebrate by contacting the DNA transcription unit in a physiologically acceptable carrier with a mucosal surface of the vertebrate.
- 8. A method of immunizing a vertebrate against an infectious agent, said method comprising administering to a mucosal (e.g., nasal) surface of a vertebrate a DNA transcription unit comprising DNA encoding a desired antigen operatively linked to a promoter region, in a physiologically acceptable carrier, thereby eliciting a humoral or cell-mediated immune response, or both, against the desired antigen, whereby the vertebrate is protected from disease caused by an infectious agent.
 - 9. A product, use or method according to any one of the preceding claims, wherein the DNA transcription unit is of nonretroviral origin.
- 25 10. Use or a method according to any one of Claims 2 to 9, wherein the antigen is viral.
 - 11. Use or a method according to Claim 10, wherein the virus is an influenza virus, e.g., virus hemagglutinin.

12. A product, use or method according to any one of the preceding claims, wherein the vertebrate is a mammal, e.g., a human.

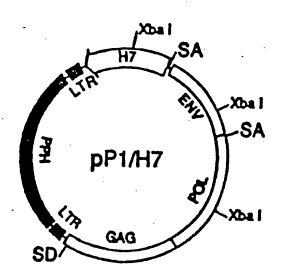


Figure 1.

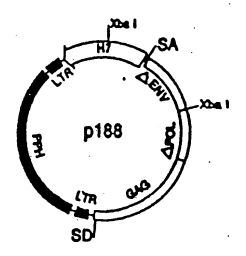


Figure 2.

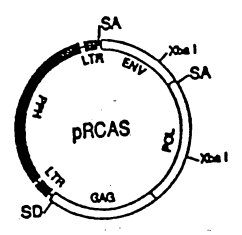


Figure 3.

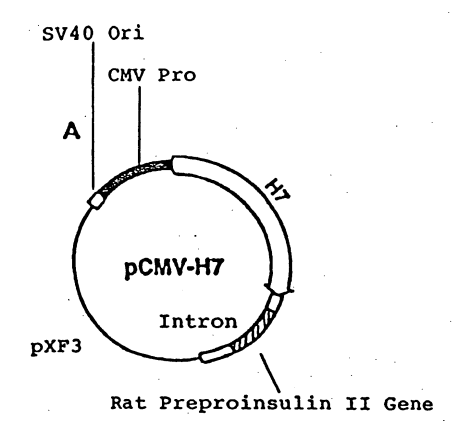


Figure 4A

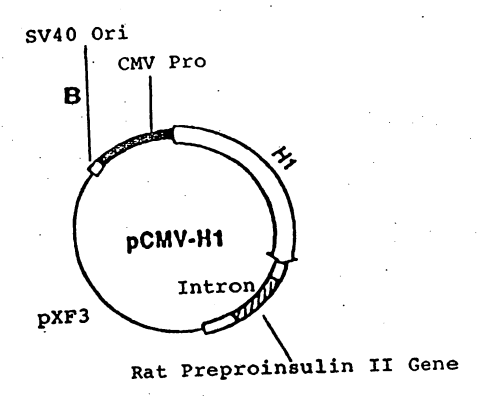


Figure 4B

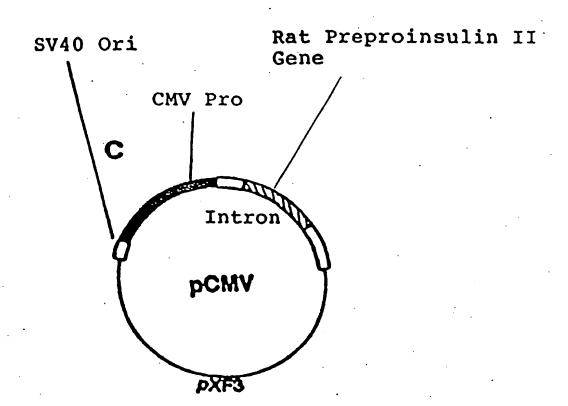


Figure 4C

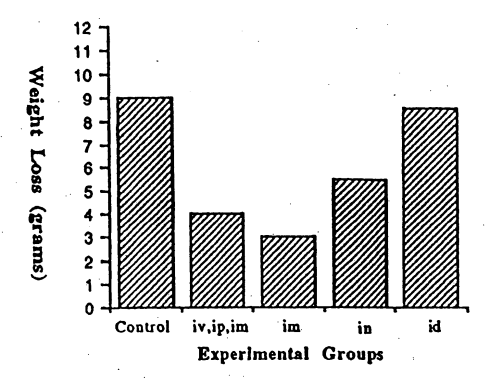


Figure 5

VIERVATIONAL SEARCH REPORT

1. CLASSIFIC	ATION OF SUBJE	CT MATTER (if several classification	symbols apply, indicate all) ⁶	
_		Classification (IPC) or to both National	Classification and IPC	
Int.Cl.	5 C12N15/4	1; C12N15/86;	A61K39/145	
II. FIELDS SI	EARCHED			
		Minimum Docum	nentation Searches ⁷	
Classification	System		Classification Symbols	
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Int.Cl.	5 .	C12N; C07K	•	
				• .
		Documentation Searched other	er than Minimum Documentation	
		to the Extent that such Documents	s are Included in the Fields Searched	
III. DOCUMI		D TO BE RELEVANT		
Category °	Citation of De	ocument, 11 with indication, where approp	riate, of the relevant passages 12	Relevant to Claim No.13
.				
X		OF VIROLOGY , no. 8, August 1988,		1-8,
) 14 - 3019		10-12
		. A. ET AL. 'Retroviru	is expressed	
		utinin protects agains		
j		za virus infections'		
		n the application		
	see the	whole document		
x	WO.A.9	011 092 (VICAL, INC.	WISCONSIN	1-6,9,12
	ALUMNI	RESEARCH FOUNDATION)		,-,
	4 Octob			
	see the	whole document		
x	WO.A.8	600 930 (WORCESTER FOU	INDATION FOR	1-6.
		ENTAL BIOLOGY)		11-12
		uary 1986		
.	see the	whole document		
	•		-/	
	•		-/	
ļ				
° Special c	categories of cited do	cuments: 10	"T" later document published after the interm	
	ment defining the ge	neral state of the art which is not	or priority date and not in conflict with t cited to understand the principle or theo	the application but ry underlying the
"E" earlie	er document but publ	Ished on or after the international	invention "X" document of particular relevance; the cit	· ·
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which	s is cited to establish on or other special r	the publication date of another	"Y" document of particular relevance; the cir	dmed invention
"O" docu	•	oral disclosure, use, exhibition or	cannot be considered to involve an invent document is combined with one or more	other such docu-
"P" docur	ment published prior	to the international filing date but	ments, such combination being obvious (in the art.	
later	than the priority dat	e claimed	"&" document member of the same patent fa	mily
IV. CERTIFI	ICATION			
Date of the A	ctual Completion of	the International Search	Date of Mailing of this International Sea	urch Report
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international (Searching Authority	A S.I. D. A STORAGE COMMON	Signature of Authorized Officer	
,	EUROPE	AN PATENT OFFICE	CHAMBONNET F.J.	
			•	

Porm PCT/ISA/210 (second sheet) (January 1965)

	International Application No						
III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)							
Category °	Citation of Document, with indication, where appropriate, of the relevant passages	Relevant to Claim No.					
	NO 4 0 COT FOR CRIOTECHNOLOGY DESEADON	1-7,10					
X	WO,A,8 607 593 (BIOTECHNOLOGY RESEARCH PARTNERS, LTD) 31 December 1986	1 7,10					
	see the whole document						
X	EP,A,O 292 879 (ORION CORPORATION LTD) 30 November 1988 see the whole document	1,9-12					
X	WO,A,9 201 045 (EQUINE VIROLOGY RESEARCH FOUNDATION, UNIVERSITY OF GLASGOW) 23 January 1992 see the whole document	1-12					
X .	WO,A,9 002 803 (INSTITUTE FOR ANIMAL HEALTH LTD, RHONE-MERIEUX SA) 22 March 1990 see the whole document	1-11					
Χ .	US,A,4 722 848 (PAOLETTI, E. & PANICALI, D.) 2 February 1988 see the whole document	1-6,9-12					
х	GB,A,2 166 349 (AMERICAN HOME PRODUCT CORPORATION) 8 May 1986 see the whole document	1-12					
X	WO,A,9 002 797 (NORTH CAROLINA STATE UNIVERSITY) 22 March 1990 see the whole document	1-6,9-11					
х	JOURNAL OF VIROLOGY vol. 64, no. 3, March 1990, pages 1070 - 1078 COSSET, F.L., ET AL. 'A new Avian Leukosis Virus-based packaging cell line that uses two separate transcomplementing helper genome' see the whole document	1					
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INTERNATIONAL SEARCH REPORT

rnational application No.

PCT/US 93/02394

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This inte	rnational search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: Remark: Although claims 3 and partially 4 to 12 as far as they concern in vivo method of treatment or vaccination against a disease are directed to a method of treatment of the human/animal body the search has been carried out and based on the alleged effects of the compound/composition.
2.	Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Roy II	Observations where with a firm in its latin (firm) and the firm is a firm of the firm of t
	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Ince	national Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2.	As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. i	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4 .	No required additional scarch fees were timely paid by the applicant. Consequently, this international scarch report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark o	The additional search fees were accompanied by the applicant's protest No protest accompanied the payment of additional search fees.

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9302394 SA 71686

This amex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information. 02/07/93

WO-A-9011092 04-10-90 WO-A-8600930 13-02-86 WO-A-8607593 31-12-86 EP-A-0292879 30-11-88	US-A- 4631191 23-12-8 AU-A- 6125386 13-01-8 EP-A- 0229826 29-07-8 US-A- 4920213 24-04-9 AU-B- 604122 06-12-9 AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8
WO-A-8600930 13-02-86 WO-A-8607593 31-12-86 EP-A-0292879 30-11-88	JP-T- 4504125 23-07-9 EP-A- 0188574 30-07-8 US-A- 4631191 23-12-8 AU-A- 6125386 13-01-8 EP-A- 0229826 29-07-8 US-A- 4920213 24-04-9 AU-B- 604122 06-12-9 AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8
WO-A-8607593 31-12-86 EP-A-0292879 30-11-88	EP-A- 0188574 30-07-8 US-A- 4631191 23-12-8 AU-A- 6125386 13-01-8 EP-A- 0229826 29-07-8 US-A- 4920213 24-04-9 AU-B- 604122 06-12-9 AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8 AU-A- 8212891 04-02-9
WO-A-8607593 31-12-86 EP-A-0292879 30-11-88	US-A- 4631191 23-12-8 AU-A- 6125386 13-01-8 EP-A- 0229826 29-07-8 US-A- 4920213 24-04-9 AU-B- 604122 06-12-9 AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8
EP-A-0292879 30-11-88	AU-A- 6125386 13-01-8 EP-A- 0229826 29-07-8 US-A- 4920213 24-04-9 AU-B- 604122 06-12-9 AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8
EP-A-0292879 30-11-88	EP-A- 0229826 29-07-8 US-A- 4920213 24-04-9 AU-B- 604122 06-12-9 AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8
	US-A- 4920213 24-04-9 AU-B- 604122 06-12-9 AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8
	AU-B- 604122 06-12-9 AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8
	AU-A- 1651488 01-12-8 CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8 AU-A- 8212891 04-02-9
	CA-A- 1300052 05-05-9 JP-A- 63304988 13-12-8 AU-A- 8212891 04-02-9
WD-A-9201045 23-01-92	JP-A- 63304988 13-12-8 AU-A- 8212891 04-02-9
W0-A-9201045 23-01-92	AU-A- 8212891 04-02-
WO-A-9201045 23-01-92	NO N
NO	
	EP-A- 0538299 28-04-
WO-A-9002803 22-03-90	AU-B- 633272 28-01-
#0 // 3002000	AU-A- 4214289 02-04-
	AU-B- 629248 01-10-
	AU-A- 4325089 02-04-
	EP-A- 0434721 03-07-
•	EP-A- 0434747 03-07- WO-A- 9002802 22-03-
•	HO 110
·	0, 1 1002000
	JP-T- 4502852 28-05-
US-A-4722848 02-02-88	8 US-A- 4603112 29-07-
•	AU-B- 561816 21-05- AU-A- 9180682 30-06-
	70 X
	E1 7130
	US-A- 5174993 29-12-
GB-A-2166349 08-05-8	
	AU-A- 4884085 08-05-
	CA-A- 1263305 28-11-
	DE-A- 3586841 24-12-

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9302394 SA 71686

This annex tists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

02/0

02/07/93

Patent document cited in search report	Publication date	Patent family member(s)		Publication date	
GB-A-2166349	-	EP-A,B JP-A- US-A-	0181117 61118326 4920209	14-05-86 05-06-86 24-04-90	_
WO-A-9002797	22-03-90	AU-A-	4307589	02-04-90	-
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more details about this annex : see (
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more details about this annex : see (Official Journal of the Euro	pean Patent Office,	No. 12/82		